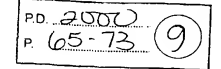
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Short Communication

DNA RESEARCH 7, 65-73 (2000)



Prediction of the Coding Sequences of Unidentified Human Genes. XVI. The Complete Sequences of 150 New cDNA Clones from Brain Which Code for Large Proteins in vitro

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Abstract

We have carried out a human cDNA sequencing project to accumulate information regarding the coding sequences of unidentified human genes. As an extension of the preceding reports, we herein present the entire sequences of 150 cDNA clones of unknown human genes, named KIAA1294 to KIAA1443, from two sets of size-fractionated human adult and fetal brain cDNA libraries. The average sizes of the inserts and corresponding open reading frames of cDNA clones analyzed here reached 4.8 kb and 2.7 kb (910 amino acid residues), respectively. From sequence similarities and protein motifs, 73 predicted gene products were functionally annotated and 97% of them were classified into the following four functional categories: cell signaling/communication, nucleic acid management, cell structure/motility and protein management. Additionally, the chromosomal loci of the genes were assigned by using human-rodent hybrid panels for those genes whose mapping data were not available in the public databases. The expression profiles of the genes were also studied in 10 human tissues, 8 brain regions, spinal cord, fetal brain and fetal liver by reverse transcription-coupled polymerase chain reaction, products of which were quantified by enzyme-linked immunosorbent assay.

Key words: large proteins; in vitro transcription/translation; cDNA sequencing; expression profile; chromosomal location; brain

We have been making efforts to accumulate information on the coding sequences of unidentified human genes.1,2 Especially, recent our interest is focused on the unidentified genes encoding large proteins in human brain since these gene products are likely to play important roles in the central nervous system. 2,3 To identify such genes, we constructed a set of strictly sizefractionated cDNA libraries from human brain and in vitro transcription/translation system have been applied to select the cDNA clones coding for large proteins prior to the determination of their entire sequence.3 As an alternative method for clone selection, we have recently introduced a computer-based approach using GeneMark analysis for picking up cDNA clones with a high probability of coding for protein.4 This new approach would be expected to minimize the risk of overlooking important cDNA clones which fail to produce proteins in vitro.

The sequences of more than 1200 cDNA clones have been reported by our project and the total length of the determined sequences exceeds 6.3 Mb¹⁻³ and the average

 Sequence Analysis and Prediction of Protein-Coding Regions in cDNA Clones

length of gene products deduced from the cDNAs from

brain is over 900 amino acid residues. 2.3 As an extension

of the preceding reports, we herein report the coding se-

quence features of 150 new cDNA clones which have the

potential to code for large proteins in vitro. In addition

sequences annotated by the database search, the expres-

sion profiles and the chromosomal locations of these 150

new genes are also described. The information regarding

these newly identified genes would greatly increase our

understanding of the biological functions of human genes

at the molecular level.

to the specific features of the newly predicted protein .

cDNA clones to be entirely sequenced were selected according to the following criteria: (1) novelties of their single-pass sequences of both the cDNA ends; (2) potentialities of their protein coding. The latter criterion was critical for us to conduct our cDNA project efficiently, because there are many cDNA clones which apparently do not possess a protein-coding region in the

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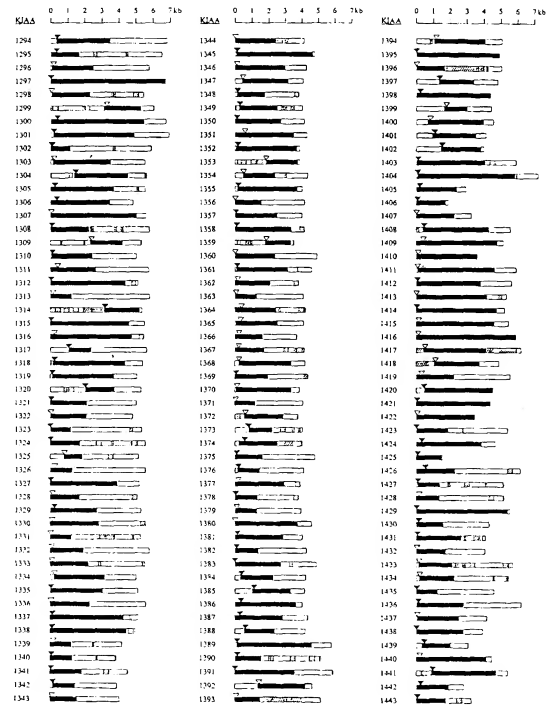


Figure 1. Physical maps of cDNA clones analyzed. The physical maps shown here were constructed from the sequence data of respective cDNA clones or, when necessary, from the combination of cDNA clones and RT-PCR products. The horizontal scale represents the cDNA length in kb, and the gene numbers corresponding to respective cDNAs are given on the left. The ORFs and untranslated regions are shown by solid and open boxes, respectively. The positions of the first ATG codons, with or without the contexts of the Kozak's rule, are indicated by solid and open triangles, respectively. RepeatMasker, a program that screens DNA sequences for interspersed repeats known to exist in mammalian genomes, was applied to detect repeat sequences in respective cDNA sequences (Smit, A.F.A. and Green, P., RepeatMasker at http://ftp.genome.washington.edu/RM/RepeatMasker.html). Short interspersed nucleotide elements (SINEs) including Alu and MIRs sequences and other repetitive sequences thus detected are represented by dotted and hatched boxes, respectively.

Table 1. Information of sequence data and chromosomal locations of the identified genes.

ene number	Accession		ORF length (amino	Chrysteemal	Gene number			ORF length (amino	Силинов
	all layers	No.	ette tengatett	he wires	(KIAA)	mers her "	Share	arid resident)"	- resipue
1294	A BAS7715	6,316	1,051	10	1340	A14137790	4,391	653	7
1295 1296	A 8037716 A 8037717	6.524 5,796	550 #15	5	1378	AR037791	LAK, E	1,107	15
1297	ARA)7718	1,796 1,726		10°	1371	A11637792	4,096	295	4
1297	AB637718	5,463	2,242 734	12.	1373	AR037793 AR037794	3,771 4,052	773 463	11 19
1199	ABM37720	KJH3	730	16	1373	48417795	4,052	164	-
1300	AR037721	6,747	1,720	15	1375	ANOJ7796	4,823	546	3,
130)	A11037722	6,926	(,34)	1	1376	AR037797	4,131	437	s'
1302	AB017723	5,904	375	H'	1377	A11837796	318.6	983	ū
1303	411037724	5,534	1,119	17"	1378	ABIL)7799	3,415	451	¥
1384	4H037725	5,433	1,051	12.	1379	A R437800	3,354	434	
1345	A15037726	5,55)	1,238	14"	13110	4 M437501	4,614	1,245	10"
1306	.11437727	4,032	1,154	16"	1381	A R837882	4,052	941	17"
1307	AB#3772#	5,601	1,67%	1	1382	A RQ37803	2112	462	13"
1208	A11037729	5,794	745	,	1383	A7637764	1,914	947	1
1304	AR437730	5,331	639	•	1384	A MAJTROS	4261	652	LA ²
1314	A H43773 I	5,028	794	2	1395	AR037806	4,193	768	14"
131\$	AH037732	5,774	XJP	5'	1384	411037807	4,030	1,214	7,
1312	AR837733	5,139	1,471	J	13#7	A 74037 NON	4742	950	2
1343	AH037734	5,314	3500	x	1364	A 863780V	20	399	16"
1217	AB437735	5,169	481	13	1389	A # 8 2 7 1 1 0	5,001	1,514	ı.
1315	.(H43773A	5,524	1,545	•	1394	A80379(1	1222	545	1
1316	48437737	5,477	1,590	11	1391	ANAJ7HI2	5,901	1,194	11,
1317	A HO 27738	5,444	435	5	1392	AB837813	4,654	950	•
אוכו לוכו	A 11037739 A 110377740	5,425 5,873	1,418 1,218	X.	1393 1394	A ROSTNIA A ROSTRIS	5,164 5,065	500 1,063	14
1326	AR437741	ינבנ	567		1395	A 8437816	4,274	1,003	12
1321	AB437742	5.054	714	17	1396	A 7437717	5,841	551	19
1323	AR437743	4,832	782	ï.	1397	A 8037718	4,819	684	6
1323	AH437744	5,356	396	18	1396	ABAJ7RIP	4.172	1,456	7
1324	AB437745	5,567	5.80	ï	1399	AB037H28	4,450	452	2
1325	A11037746	5,155	354	4	1406	AH037#11	4.554	1,093	4
1324	A HO.17747	5,34,1	424	14	1401	AHAJ7R22	4.187	LCS	17
1327	A H0 377 44	5,205	1714	4'	1402	A 7637813	3.979	788	17"
1328	ARE37749	3,047	\$74	15	1463**	AR037#24	5,497	1,137	15
1.329	A36437750	5,2117	967	4	1404	A RO37#25	7,204	1,525	20
13.18	AB437751	5.577	945	15	1405	A 8837426	2,545	751	1
1331	A19437752	5,273	412	٦,	1406**	AB037#27	1,176	571	,
1333	A 11037753	5,71s	631	C,	1407	AH037H28	3247	744	3,
1333	A 11#37754	5,534	741	14	1401	AROJ7K29	5,548	1,298	19*
1334	A H#37755	5,643	7173	S .	14997	A11037#30	5,140	1421	14
1,335	4 84 37756	5.123	1,026	26	14100	A #037#31	3,644	1,291	1
1336	A 84137757 A 8413775%	5,591 5,181	766 1,438	1 ⁻	1411 ^{et} 141 <i>2</i> et	ARG37833 ARG37833	5,91RI 1.34.7	L322 L374	6°
1337	AB037759	4,994	1,495	15.	1417	ARH37834	3,261 3,261	1,274	,
1774	5 Ro3776#	4.217	409	7	1414	AB037835	5,242	1,5%	;
1346	A 8617761	3.476	41	12	1415**	AIM37834	5.450	1,539	20
1341	A11837742	4,544	620	15"	1416"	AIIAI7AI7	5.901	1.367	R
1.42	A 80,5776,3	3,210	426	18"	1417*	A199374.48	6,204	1,217	,
1343	A BA 3776-6	440	520	i.	1418**	A BA37839	4,396	AL 9	1.
1344	AB037745	4.135	106	14"	1419**	A R437448	5.540	738	ū'
1745	ABBUTTES	4,750	1,532	4"	1428"	AMOTALL	4,516	1,39	3
1346	4 RH37747	4_309	999	21	1421"	A1037#42	4,391	1,443	15
1,347	A H037764	4,073	918	3	1422	AIM37#43	3,456	1,151	,
1344	A PALIT769	3,410	515	16	142,5**	ATI 137444	5,340	416	4.
1.49	AB037770	4,1155	752	17	14241	A 219 37 845	4,655	1,2%	4
1334	AIMLITTT I	4,153	911	4.	1435***	A 11037446	1.543	495	٠,٠
1351	AB037772	4_347	1,163	10,	1426*	4 8437 447	6,140	758	16
1352	4 BB37773	3,593	1,213	5'	1427#	A110.37%48	5,145	439	11,
1353	A R037774	3,477	418	1	HUP	ANDITAR	5,148	458	3
1354	A RUST/75	4,352	431	•	1-1290	ABBJ7850	5,507	1,795	4,
1355	A8037776 A8037777	4,036	1,159	•	14345	AD837851	4,282	527	4
1354	A11937777	4,193 4,022	519 834	1 6	1431°*	A 88 37 8 52	4,076	791	19
1354	A 11937777	4,122 4,133	1.123	7	1432"	AR037853 AR037854	4.02.1 5.67.1	571 452	•,
1359	A H& 577#U	4,183 3,554	1.123 517	, ,	1433**	47437454 48437855	5,443	677	3.
1359	A HO 3 7 7 M L	3234	794	12`	1435°	4 R037R33	3,443 4,574	415	20°
1.341	A 88377#2	4,625	1,005	17.	1436*	ABO 17857	6,160	924	ŕ
1342	A ##377#3	3,542	499	12	14375	AH037854	4.161	7 <i>(</i> 4	· ·
1343	ABOJT7 W	4116	14)	ĵ.	1437	ADU37859	1,907	734	72
1364	AHAJ7715	4,261	411	21	(4)	A 70 37 860	J,043	541	ii.
1365	A 18,17786	4,150	101	ï	14000	A 8037861	4,434	777	,
t Jea	4 8037757	3,714	554	17*	1441'*	48037962	5,37A	1,254	è
1347	4 H4 177 Rd	4,196	579	14"	(442**	ARAITRAL	2,7402	627	19
1,368	AR437719	4.250	1.047	5.	144314	AB037XA-I	3,214	573	14

a) Accession numbers of DDBJ, EMBL and GenBank databases. b) Values excluding poly(A) sequences. c) Values were calculated from the number of amino acid residues between two termination codons in the case where the in-frame termination codon exists upstream of the first ATG codon. d) Chromosome numbers were identified by using GeneBridge 4 radiation hybrid panel unless specified. The actual primer sequences and the PCR conditions used for the radiation hybrid mapping are accessible through the World Wide Web at http://www.kazusa.or.jp/huge. The chromosomal locations highlighted by asterisks were fetched from the UniGene database. The chromosomal locations highlighted by sharp were referred from the GenBank database because the sequences of the cDNA clones could be found in the genomic sequences whose chromosome numbers were assigned. e) cDNA and ORF lengths were revised by direct analysis of the RT-PCR products. f) Nucleotide sequences were determined after subcloning of the internal Not I-digested fragment. Therefore, cDNA length of these genes represented those of internal Not I-digested fragment. g) cDNA clones were selected by analysis of 5'-end single-pass sequences using the GeneMark analysis.

Table 2. Functional classifications of the gene products.

2-1. Predicted function based on homology search^{a)}

Function*	Gene product	44 TS.	OWL ID	475 4	identity 9		
Cell signaling/communication	KIAA1296	815		714	82	COVERIE.	Definition
	KIAA1297	22.42		1431	35	96	ports in-1, complete cds mouse
	KIAA1299	730	JC5887	670	93	13	
	KIAA1304	i 05 I	P98171	946	48	92	signaling mediator variant - mouse
	KIAA1308	745	O03385	852		72	
	KJAA1312	1471	D67C76		81	73	guantite nucleoridedissociation atimulator in ICDS form A
	KIAA1314	681	Y00661	951	44	46	secretory protein containing thrombospondin mouls, complete cds mou
	KIAA1322	702	UB1500	1227	30	30	bet - hartian
	KIAA1327	1310	T03730	438	39	50	phgA gene, complete eds Dicryostelium discoideum
	KJAA1338	1495	M20487	1567	6;	100	antigen containing epicope to monoclonal antibody MMS-85/12 - mouse
	KJAA1342	426		1020	35	31	protein kinase GCN2, complete cds 5 cerevisine
	KJAA 1347	918	P50232	425	90	100	synaprotagmin IV - rat
	XIAA1348		A 42764	919	97	100	Ca2+transporting ATPase (EC 3.6.1.38) - rat
		545	AF062741	530	14	97	Territoria della America della J.O. (1.36) - Tel
	KIAA 1356	519	P08104	1951	97	100	pyruvate dehydrogenase phosphatase isoenzyme 2, complete cds rai
	KJAA1361	1005	AF084205	1001	99	100	sodium channel protein, brain I alpha subunit - human
	KJAA 1366	550	U41662	836	98	100	serime/threonine protein kirase TAOI, complete cds rat
	KCAA1368	1049	AF030430	388	93	84	neuroligin 2, complete cds rui
	KIAA 1369	653	AF028808	619	15		scrnaphorin VIa, complete cds mouse
	KIAA 1385	768	Q03555	736	100	95	hemin-sensitive initiation factor 2 alpha kinase, complete eds mouse
	KIAA1389	1514	AF090989	1783	53	96	gepriytin (putative glycine receptor-tablic linker nove in) . ex-
	KIAA 1400	1093	U88549	896	97	96	putative GAP protein alpha, complete eds human
	KIAA 1422	1151	AF089730	1237		3C	OL-protocacherin, complete cds. + mouse
	XIAA 1424	1286	U02289	1439	94	91	pocussium channel subunit (Sleek), complete eds - rat
	KIAA 1427	439	P46096		48	17	GTPase-activating protein (CEGAP), partial cds C elegant
	KIAA1436	924	O62786T	421	32	61	Symaprotagmin i - mouse
or lese acid management	KIAA1339	409	AF020591	879	89	95	prostaglandin F2-alpha receptor regulatory procein precursor - rat
	KIAA1341	620	A56704	715	45	61	zinc finger protein, complete cds - human
	KIAA1349	752		435	90	73	regulatory protein Myef-2 - mouse
	FJAA1367		Q05481	1191	56	38	zinc furger protein 43 - human
	KIAA1380	579	Q10368	782	99	100	Cleaner and and a day had
		1265	Q63679	1714	46	66	cleavage and polyadenylation specificity factor, 100 kD subunit - bovine
	KIAA1388	599	Q05481	1191	39	83	testis specific protein A - rac
	MAA1396	551	P52742	469	59	83	zinc finger protein 91 - human
	KIAA 1416	1967	X86691	1912	42	83 34	zinc finger protein 135 - human
	KIAA1431	89 i	P10078	614	75	64	218kD Mi-2 - human
	KIAA1439	561	P09414	509	100		zinc finger protein ZFP28 - mouse
	KIAA1442	627	U92704	55;	77	91	nuclear factor 1 (NF-I) - rat
Otean management	KJAA1443	573	JC4863	873		83	Olf-I/EBF-like-2(OS) transcription factor, complete cds mouse
Otem management	KIAA1.01	1581	P46934	927	35	35	homeonic protein protein zhzil - mouse
	KIAA1320	567	AF037454	854	36	49	KIAA0322, pertial cds human
	KIAA1346	999	T000:7	951	45	61	ubiquitin protein ligase, complete cds mouse
	KJAA1352	1212	009996		62	95	ADAMTS-1 protein - mouse
cubolism	KIAA1363	430	A58922	1;98	56	97	probable leucyl-IRNA synthesase (EC 6 1.1.4) - C. elegans
Il structure/motility	KIAA1294	1051	P26044	398	43	94	esterate N-descetylase (EC 3.5.1-), 50K hepatic - rabbit
	KIAA1306			583	32	24	radium - pig
	FJAA1309	1154	572647	464	35	18	extensin - Volvoz corteri
		639	AF059569	593	30	85	
	KFAA1354	632	AFC59569	593	30	86	actin binding protein MAYVEN, complete cds human
	KIAA1357	836	\$21697	464	ĴŠ	25	actin binding protein MAYVEN, complete cds human
	XIAA1362	699	AF038388	766	33		extensin - Volvos carreri
	KIAA1365	831	U66707	1495		.64	actus-filament binding protein Frabin, complete cds rat
	ETAA1578	451	AF059559	593	93	:00	densin-180, complete eds rat
	KIAA1405	791	AF009624	393 742	36	95	actin binding protein MAYVEN, complete eds human
	F14A1410	1201	U03975		91	30	K1F3-related motor protein partial cds - human
	KJ 4 A T 4 3 7	817	U667C7	1125	77	68	dyneur heavy chain isotype 6, partial cds sea urchin
		911	000767	1495	30	38	densin-180 complete cds rai

a) Homology search was performed by Smith-Waterman algorithm, using BioView Toolkit and GeneMatcher (revision 3.3, Paracel Inc. USA) against OWL database (release 31.4). The homologous protein with the highest score was listed, when it satisfied the following conditions, i) the protein was functionally annotated, ii) the aligned region exceeded 200 amino acid residues, and iii) percent identity in the algined region was 30% or greater. b) Function was classified based on the annotation of the entry of the homologous protein in the database. c) The values mean the ratio of the length of aligned region to the original length of the query sequence, in percentage.

cDNA libraries derived from tissue poly(A)⁺ RNA. To screen cDNA clones according to their protein-coding capability, we have used an *in vitro* expression system and recently introduced a computer-based method called GeneMark analysis for minimizing the risk of overlooking important cDNA clones.^{2,4} In this report, 21 cDNA clones were selected by GeneMark analysis and 129 cDNA clones were selected by the *in vitro* expression system. These cDNA clones were isolated from the size-fractionated human adult brain cDNA libraries Nos. 2 to 5 (insert sizes ranging from 4 to 6 kb) and the size-fractionated human fetal brain cDNA libraries Nos. 4 and 6 (insert sizes ranging from 4 to 7 kb) previously constructed.^{2,3} The clones with unidentified sequences at both ends were chosen by single-

pass sequencing and a homology search was performed against the GenBank database (release 113.0) excluding expressed sequence tags and genomic sequences.³ A total of 35 cDNA clones (KIAA1389-KIAA1402, KIAA1415-KIAA1422, KIAA1424, KIAA1425 and KIAA1433-KIAA1443) were selected from the adult brain libraries and the remaining 115 cDNA clones were obtained from the fetal brain cDNA libraries. Entire sequencing of these clones was performed according to the methods previously described in detail.^{2,3} Twenty-three clones (KIAA1403-KIAA1425) seemed to carry spurious coding interruption caused by errors of the reverse transcriptase or by retained intron sequences. For these cases, the sequences of the regions causing interruption of an open reading frame (ORF) were reexamined by direct se-

Table 2. Continued.

2-2. Predicted function by motif search^{a)}

Function**	Gene product	aa res.	Pfam ID	E-value"	Definition
Cell signaling/communication	KIAA1295	550	PF00018	4.30E-06	SH3 domain
			PF00018	1.30E-04	SH3 domain
	KIAA1298	738	PF00782	2.10E-34	Dual specificity phosphatase, catalytic domain
	KIAA1330	945	PF00047	4.10E-02	Immunoglobulin domain
	KIAA1355	1189	PF00041	1.50E-09	Fibronectin type III domain
		,	PF00041	1.80E-08	
			PF00047	5.70E-01	Fibronectin type III domain
			PF00047		Immunoglobulin domain
			PF00047	4.20E-12	Immunoglobulin domain
			PF00047	3.60E-08	Immunoglobulin domain
			PF00047	5.50E-05	Immunoglobulin domain
	K[AA1360	704		9.00E-06	Immunoglobulin domain
	KIAA1391	796	PF00069	3.00E-07	Eukaryotic protein kinase domain
	VIVA1391	1194	PF00169	9.30E-01	PH domain
	1711 11107		PF00620	7.30E-30	RhoGAP domain
	KIAA1406	1876	PF00888	4.00E-01	Cullin family
	KIAA1415	1539	PF00610	1.70E-10	Domain found in Dishevelled, Egl-10, and Pleckstrin
	KIAA 1428	458	PF00169	5.10E-04	PH domain
Maria and a			PF00640	3.70E-04	Phosphotyrosine interaction domain
Nucleic acid management	KIAAI3II	389	PF00075	5.90E-02	RNA recognition motif
			PF00642	3.50E-02	Zinc finger C-x8-C-x5-C-x3-H type
	KIAA1343	520	PF00249	1.80E-08	Myb-like DNA-binding domain
			PF00249	4.10E-06	Myb-like DNA-binding domain
			PF01448	3.30E-12	ELM2 domain
	KIAA1384	652	PF00651	2.60E-24	BTB/POZ domain
			PF01344	4.10E-02	Kelch motif
			PF01344	7.60E-03	Kelch morif
			PF01344	5.10E-15	Kelch motif
			PF01344	5.20E-06	Kelch motif
			PF01344	5.90E-05	Kelch motif
			PF01344	1.20E-01	Kelch motif
	KIAA1425	495	PF00249	9.20E-01	Myb-like DNA-binding domain
	KIAA1441	1258	PF00096	3.10E-02	Zinc finger, C2H2 type
			PF00096	6.50E-02	Zinc finger, C2H2 type
			PF00096	9.80E-04	Zinc finger, C2H2 type
			PF00096	2.30E-02	Zinc finger, C2H2 type
			PF00096	5.20E-03	Zinc finger, C2H2 type
ell structure/motility	KIAA1364	811	PF00307	8.60E-18	Calponin homology (CH) domain
			PF00412	3.30E-06	LIM domain containing proteins
rotein management	KIAA1333	741	PF00632	2.20E-01	HECT-domain
	KIAA1350	911	PF00443	6.30E-01	Ubiquitin carboxyl-terminal hydrolase family 2
	KIAA 1372	773	PF00442	4.10E-13	Ubiquitin carboxyl-terminal hydrolases family 2
			PF00443	9.10E-20	Ubiquitin carbonal terminal hydrolases (amily 2
	KIAA1414	1586	PF00298	1.40E-01	Ubiquitin carboxyl-terminal hydrolase family 2 Ribosomal protein L11
fetabolism	KLAA1315	1545	PF00389		N10030III4I pi016III LTI

a) Motif search was performed by HMMER2.1.1 against Pfam database (release 4.4). b) Function was classified based on the annotation of the Pfam entry which was hit in the query sequence. c) Only the entries possessing the expectation value (E-value) less than 1.0 were presented.

quencing of the major reverse transcription-coupled polymerase chain reaction (RT-PCR) products to precisely predict protein-coding sequences.⁵ This examination revealed spurious interruptions in the following clones: ORFs in 7 clones (KIAA1403, KIAA1405, KIAA1409, KIAA1410, KIAA1415, KIAA1424 and KIAA1425) were found to carry single- or multiple-insertions most of which probably corresponded to intronic sequences; ORFs in 7 clones (KIAA1411, KIAA1412, KIAA1413, KIAA1416, KIAA1418, KIAA1420 and KIAA1421) were frame-shifted by single- or double-short insertions or single-deletion (< 5 nucleotide residues); ORFs in 4 clones (KIAA1404, KIAA1408, KIAA1417 and KIAA1423) were found to carry single- or doubledeletions; ORFs in 4 clones (KIAA1406, KIAA1407, KIAA1414 and KIAA1422) were divided into some por-

tions by a combination of spurious interruptions including insertions/deletions; KIAA1419 carried a nonsense mutation in the ORF. For those genes, the revised sequences by the RT-PCR experiments, not the actual cloned cDNA sequences, were deposited to Gen-Bank/EMBL/DDBJ databases and used for analyses in this study including prediction of their protein-coding sequences unless otherwise stated. The results of the comparison between the cloned DNA and the revised DNA sequences are available through the World Wide Web site at http://www.kazusa.or.jp/huge. The actual primer sequences and the PCR conditions used for the RT-PCR experiment are accessible through the web site http://www.kazusa.or.jp/~hirosawa/interruption/ entrance.html. Notably, clones for eight genes (KIAA1297, KIAA1398, KIAA 1395, KIAA1410,

Table 3. Homologues of the newly identified genes found in various databases. a)

Danhaa."	New gene	B. 51.	ID in database	12.771	% karany "	h,m,rege*	Comnent
HI'GE AND INTO ALTER	KIAA1294	1051	KIAA (O13	1062	31	90	
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	KIAALMI	1581	KIAA0122	1562	36	98	
	KIA A I 304	1051	KIA AO456	1095	68	99	
	KIAA 1306	1154	KIAA 1139	1124	34	100	
	KIAA1309	634	KIAA1354	631	92	93	
		639	KIAA1129	625	30	86	
	KIAA 1116	1590	KIAA1414	15346	36	98	
	KIAA 1146	999	KIAA0688	849	49	RI	
	KIAA 1347	915 751	K1A A0703	1051	45	96	
	KIAAT 149 KIAAT 154		KIAA1141 KIAA1129	625	30	86	
	KIAAT 161	632	KIAAOREI	1064	70	100	
	KIAAL166		KIAA0951	679	61	100	
	KIAA I 178	451	K1AA0795	465	15	96	•
	SWIALS	551	KIAA0796	682	50	#.X	
	KIAA1431	791	KIAA0065	148	41	86	(
	KIAA1441	1258	KIAAOZII	1317	34	92	
C24	KIAA1347	918	SW ATCI_YEAST	950	30	93	Cu2+4renuporling ATPase (EC 3.6.1.38)
	KIAA 1152	1212	SW-SYLC_YEAST	1090	46	84	lencyl-IRNA synth, tasc, cytoplasmic (EC & 1.1.4)
	_KIAA1401	E53	\$47545	722		90	Impathetical progrim YDC0600
degnas	KIAA 1 147	918	ZK254.1x	923	.59	94	CECC42
			K11D9.26	1004	34	81	CEXTIDAL
			K1109.25	1059	36	8.6	CEX11D92
			B0345.2	996	30	15	CEB1/652
			C01C12.8	1049	30	92	CECOIG 126
	KIAA 1352	1212	R74.1	1186	56 37	47	probable leucyf-IRNA xyrchitase (EC 6.1.1.4)
	KIAA 1361 KIAA 1374	1005 764	717E9 (F3AG1.)	982 759	41	29 95	stone/hieronine protein kinase satu (EC 2.7 1) the 2 materia
	KIAA1374	/64 431	R1252.1	531	39	בר ונ	CELR12E214
	KIAATATI	431 853	FIOC7.1	785	.77 39	• 1	CELF/0G79
	KIAA1422	1151	PORB 12.3%	1107	46	83	CEP08122
	71771742	****	FDARIZ In	1119	46	Ñ	CFR(8)71
	KIAA1434	677	TOSH 10.7	796	33	90	hyputhrical 90.8 kd protein iOSh10.7 in chromosome []
	Alan, 199	•.,	K1083.6	757	30	93	CPLX (08.1)
	KIAATAN	413	02013 2	415	40	95	hypothetical 46.2 hd up-asp retems contaming prospin (20) 3.2 is chromosome II
OW1	KIAA1296	315	APOTM667	714	87	96	strate), complete cita masur
-	KIAA1299	730	JC5847	670	91	92	Migneling mediator verters - mouse
	KIAA I 101	1581	KIAA0322	1562	54	94	KIAA0322, partial cds horson
	KIAAIJQI	1119	SPAC57A710	(11)	18	98	S.pombe chronsumme I countid c57A7 franch years
	KIAA 1304	1051	KTAADASA	1002	ė#.	99	KIAA0456, partial culs Numm
	KIAALYO	619	A PO39569	593	30	85	action trinking protein MAYVEN, complete cube - burners
	KIAA:327	1910	70,1730	1567	41	100	antigen creatining epitope to monaclonal arcibody MMS-RS/12 - mraar
	KIAA 341	620	\$15537	729	45	87	hrRHA-binding protein M4 - human
	KIAA1342	426	SYTA_RAT	425	90	100	Symphologram (V - cul
	KIA A 1.146	999	TINIT	951	#2	95	ADAMTS-1 protein - re-more
	KIAALSIT	912	A 42764	919	97	(AD)	Ca2+-transporting ATPace (EC 1.6.1.34) - net
	KIAA134# KIAA1349	345 752	A P062741	3 10 803	114 54	97	pyravise dehydrogenaat phosphatair fatericyme 2, complete eds hit
	KIAA1357	1212	ZN4J_HUMAN	1198	54 54		sinc finger process 43 - human
	KIAA1 134		SYLC_CAEEL ARISYS69	593	36 30	97	probablicacyticRNA symbolian (EC &1.1.4) - C. elezano
	K1AA £356	&12 319	CTN I_HUHAN	42)	91	# I	actio hinding protein MAYVEN, complete cita - buntum malium changed protein, brain f alpha nationa - buntum
	KIAA: MI	1005	AFREAZOS	3001	99	100	sense therein, react a process to the sense to the control of the
	KIAALJAT	410	A 5#922	198	43	94	entertac/N-de sectylase (EC 3.5.1-), 50K hepatic - rabbit
	KIAAIIM	1049	APCVIANG	444	91		ALMONIAN AIT COLLEGE AND A STATE AND
	KIAA1169	653	A PITZ NIBOR	619	x.3	V5	heren servicine installan factor 2 alpha kinner, complete cult monte
	KIA 41.173	461	HSU73522	424	37	87	AMSH, cumplest rule human
	KIAA1174	264	CEL011523	740	41	98	CHE 2 property - C. physical
	KIAA: 176	437	\$735VIB	391	41	19	brain expressed HHCPA78 humolog - human
	KIAALITE	451	KIAAU795	463	35	94	KIAA0795, perual cde hernen
	KIAA1379	434	AF104412	441	96	Inc	gredages 1, complete cula - rat
	KIAA (181	94.1	AF109377	980	#2	99	LFBp (LDLB), complex cds move:
	K1A A13*2	462	H3U+90#3	304	37	DR	transporter protein (g.17), complete cdu human
	KIAA;385	768	CEPH_RAT	736	100	944	pophysin (paget) ve plycone receptor tublis linker present) - rm
	KIAAINER	199	ZIM_HUMAN	724	3.5	F2	son: finger pusein 184 - human
	VEE I A A F 2	1514	A PONOVA 9	1783	53	96	parative GAP protein alpha, complete cula - homan
	KIAA1191	300	JC4255	475	33	#2	recs-10+ protein - Meurosperia crissia
-	KIAAL196	551	ZIJS_HUMAN	469	39	- 83	zinc Einger protein 135 - human
	KJA A 1398	1456	A 56714	1534	KI .	93	obsione receptor, 190k - dog
	KIAAI#IQ	11791	MMC88349	296	97	R 0	OU-protections, complete cult. mount
	KIAA 1411 KIAA 1422	853	CELP 10G79 APUR9730	785	39	91	Ca. norhabitis elegans cosmid F10C7 - C. elegans
		1131	ZIII-HUMAN	1237	**	91	principle charged suburus (Slack), complete colo. Int
	KIAA1431 KIAA1413	652	ZIIII_HUMAN APOSITAR	726 630	45 39	# 3 9-4	ranc Einger pratein 184 - harmon
	KIAA1434	677	YRS7_CAEEL	796	31	90	hrain specific contactin himbrig protein CBP90, partial cds rat hypothetical 90,8 KD protein T03H10.7 in chromosome II - C. elegans
	X1441435	415	YLN2_CAEEL		,ii	*0	
	KIAA1416	924	FPRP RAT	415 279	40) 10	95	hypichicical 46.2 kd trp-aup reprote containing protein (2013.2 in chromosome 11 - C. elegent
	KIAA1419	923 561	MFTL_RAT	309	100	95	prottaglandin F3-alpha receptor regulatory pristrin procursor - rat nuclear factor ((NF-I) - rat
	XIAA1441	1238	DA6966	1267	100 24	91	KIAADIII. complete cds human

a) The definition of homologues used here was the proteins found in the databases satisfying the following conditions: i) the length ranged from 80% to 125% of the query sequence; ii) the ratio of the length of aligned region to that of the original sequence of the query was 80% or greater; iii) percent identity was 30% or greater. The method of homology search was the same to that explained in Table 2-1. b) The following databases were used. HUGE, our cDNA-encoded protein database (http://www.kazusa.or.jp/huge); yeast, non redundant peptide database from genome-ftp.stanford.edu:/pub/yeast/yeast_protein/yeast_nrpep.fasta.Z; C. elegans, protein database deduced from C. elegans full genome sequence (ftp.sanger.ac.uk:/pub/databases/C.elegans_sequences/C_elegans_proteins_1998-10-16.pep) and the entries derived from C. elegans of OWL, and OWL (release 31.4). In the case of database search against OWL, only the homologue with the highest score to each query was listed. c) The number of amino acid residues of the gene produt. d) The values atean the ratio of the length of aligned region to the original length of the query sequence, in percentage. e) For entries from databases, yeast and OWL, the annotations were listed. For C. elegans, IDs of OWL were listed, when sequences identical to the entries from the full genome were registered in OWL.

KIAA1416, KIAA1420, KIAA1421 and KIAA1422) seemed to lack regions encoding C-terminal portions due to the presence of a Not I site in their coding regions because cDNAs were digested with Not I before ligation into vector. In contrast, clones for five genes (KIAA1439-KIAA1443) were found to lack 5'-portions of the sequences due to the presence of an internal Not I site in their sequences. For these five genes, the nucleotide sequences of only the region between two NotI sites were determined, since their original clones were most likely to harbor two intermolecularly ligated independent cDNAs.6 After these revisions, the average size of the cDNA sequences became 4.8 kb and that of the ORFs corresponded to approximately 910 amino acid residues. Physical maps of the 150 cDNA sequences analyzed are shown in Fig. 1, where the ORFs and the first ATG codons in respective ORFs are indicated by solid boxes and triangles, respectively. Repeat sequences are also shown in Fig. 1. Comparing the predicted proteincoding sequence for KIAA1299 with those of mouse and rat homologues, 7.8 this cDNA clone seems to encode a complete protein although it possessed an unusually long 5' non-coding sequence expanding more than 3 kb. Table 1 lists the lengths of inserts, the ORF lengths and the chromosomal locations of the respective clones. Chromosomal loci of 66 newly identified genes were assigned using human-rodent hybrid panels, GeneBridge 4 (Research Genetics Inc., USA), since their mapping data were not available in the public databases. The chromosomal locations of the 78 genes, which are highlighted by asterisks in Table 1, were fetched from the UniGene database (http://www.ncbi.nlm.nih.gov/UniGene). The chromosomal locations of the remaining six genes, which are highlighted in Table 1, were obtained from the Gen-Bank database because the sequences of the cDNA clones were already assigned to chromosome numbers.

2. Functional Classification of Predicted Gene Products

The gene products predicted from the cDNA sequences were classified by homology and/or motif search against the following public databases: protein sequence database, OWL (release 31.4), 10 databases of predicted protein sequences from yeast 11 and C. elegans 12 genomes [genome-ftp.stanford.edu:/pub/yeast/yeast_protein/ yeast_nrpep.fasta.Z, ftp.sanger.ac.uk:/pub/databases/C. elegans_sequences/C_elegans_proteins_1998-10-16.pep!, protein domain database, Pfam (release 4.4), 13 and our own database, HUGE14 (http://www.kazusa.or.jp/ huge). As shown in Table 2, the 73 gene products were classified into five functional categories. Among them, 53 gene products indicated significant sequence similarity to functionally annotated proteins (Table 2-1). The functions of the other 20 gene products were predicted based on the presence of functional motifs/domains,

since they did not show sequence similarity to functionally annotated proteins (Table 2-2). In total, 63 gene products (86.3% of genes functionally annotated here) were suggested to have functions relating to cell signaling/communication, nucleic acid management or cell structure/motility. Of the 12 genes in functional class of nucleic acid management, 5 coded for DNA binding proteins carrying C2H2-type zinc finger domains. The average number of these domains among these gene products was about 15. Since the majority of zinc finger proteins in yeast contain only two domains per polypeptide, multiple appearance of C2H2-type zinc finger domains in a single polypeptide might be a specific character of large proteins in multicellular organisms. To find the genes conserved in other species, we tentatively defined "homologues" as genes sharing at least 30% of protein sequence identity spanning almost the entire region (more than 80% coverage against the query protein sequence). As shown in Table 3, 48 KIAA gene products were found to have the "homologues" in the databases. Homologues to 9 of the 48 KIAA proteins were found in C. elegans and 3 (KIAA1347, KIAA1352 and KIAA1401) were found in both yeast and C. elegans.. KIAA1347 and KIAA1352 were similar to Ca2+-transporting ATPase and leucyltRNA synthetase, respectively, though KIAA1401 had no similarity to any functionally known genes.

3. Expression Profiles of Predicted Genes

The expression profiles of the genes newly identified in this study are shown in Fig. 2 by using color codes. ¹⁵ KIAA1379 was homologous to rat synaptic dynamin-associated protein I (Syndapin I)¹⁶ and predominantly expressed in hippocampus. The gene expression levels of KIAA1341 and KIAA1366, which were similar to mouse transcriptional suppressor of the myelin basic protein gene¹⁷ and rat neuroligin 2, ¹⁸ respectively, were relatively high in all brain regions examined. KIAA1346 and KIAA1434 were predominantly expressed in spinal cord. KIAA1312, KIAA11315 and KIAA1417 were expressed very poorly in all regions examined, but their mRNAs were detected. These expression profiles also provide us important information for identifying biologically important genes characterized in this project.

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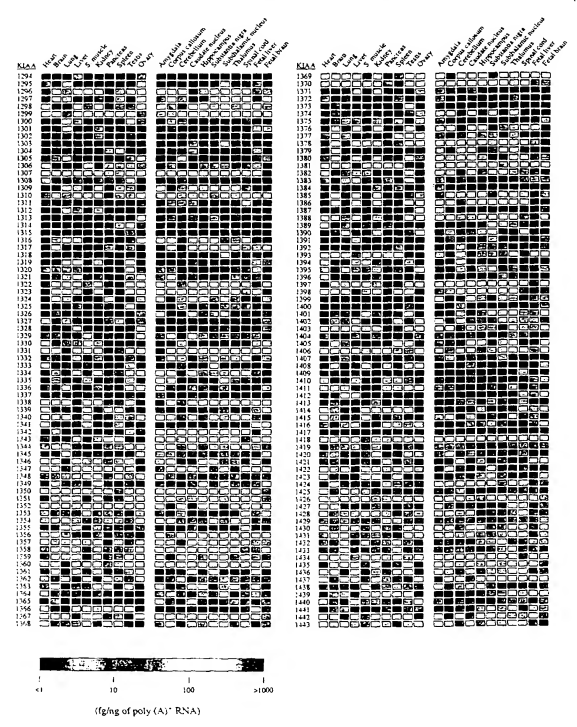


Figure 2 Expression profiles of 150 newly identified genes examined by RT-PCR ELISA. The tissue expression levels of the 150 human genes were analyzed by using the RT-PCR ELISA according to methods previously described. Gene names are given as KIAA numbers at the left side of each set of color codes. Tissue and brain region names are indicated above the top sets of color codes. A color conversion panel shown at the bottom was used for displaying mRNA levels as color codes. The mRNA levels are expressed in equivalent amounts (fg) of the authentic cDNA plasmids in 1 ng of starting poly(A) RNAs. Besides 10 tissues, 9 regions of the adult central nervous system (amygdala, corpus callosum, cerebellum, caudate nucleus, hippocampus, substantia nigra, subthalamic nucleus, thalamus, and spinal cord) and fetal brain were included in the expression profiling. As a control, mRNA levels in fetal liver were also examined.

References

- Nomura, N., Miyajima, N., Sazuka, T. et al. 1994, Prediction of the coding sequences of unidentified human genes. I. The coding sequences of 40 new genes (KIAA0001-KIAA0040) deduced by analysis of randomly sampled cDNA clones from human immature myeloid cell line KG-1, DNA Res., 1, 27-35.
- Nagase, T., Ishikawa, K.-I., Kikuno, R. et al. 1999, Prediction of the coding sequences of unidentified human genes. XV. The complete sequences of 100 new cDNA clones from brain which code for large proteins in vitro, DNA Res., 6, 337-345.
- Ohara, O., Nagase, T., Ishikawa, K.-I. et al. 1997, Construction and characterization of human brain cDNA libraries suitable for analysis of cDNA clones encoding relatively large proteins, DNA Res., 4, 53-59.
- Hirosawa, M., Nagase, T., Ishikawa, K.-I., Kikuno, R., Nomura, N., and Ohara, O. 1999, Characterization of cDNA clones selected by the GeneMark analysis from size-fractionated cDNA libraries from human brain, DNA Res., 6, 329-336.
- Ishikawa, K.-I., Nagase, T., Nakajima, D. et al. 1997, Prediction of the coding sequences of unidentified human genes. VIII. 78 new cDNA clones from brain which code for large proteins in vitro, DNA Res., 4, 307-313.
- Nagase, T., Ishikawa, K.-I., Miyajima, N. et al. 1998, Prediction of the coding sequences of unidentified human genes. IX. 100 new cDNA clones from brain which can code for large proteins in vitro, DNA Res., 5, 31-39.
- Riedel, H., Wang, J., Hansen, H., and Yousaf, N. 1997, PSM, an insulin-dependent, pro-rich, PH, SH2 domain containing partner of the insulin receptor, J. Biochem., 122, 1105-1113.
- 8. Rui, L., Mathews, L. S., Hotta, K., Gustafson, T. A., and Carter-Su, C. 1997, Identification of SH2-Bbeta as a

- substrate of the tyrosine kinase JAK2 involved in growth hormone signaling, Mol. Cell Biol., 17, 6633-6644.
- 9. Gyapay, G., Schmitt, K., Fizames, C. et al. 1996, A radiation hybrid map of the human genome, Hum. Mol. Genet., 5, 339-346.
- Bleasby, A. J., Akrigg, D., and Attwood, T. K. 1994, OWL - a non-redundant composite protein sequence database, Nucleic Acids Res., 22, 3574-3577.
- 11. Goffeau, A., Barrell, B. G., Bussey, H. et al. 1996, Life with 6000 genes, *Science*, 274, 546-567.
- 12. The C. elegans Sequencing Consortium. 1998, Genome sequence of the nematode, C. elegans: A platform for investing biology, Science, 282, 2012-2018.
- Bateman, A., Birney, E., Durbin, R. et al. 1999, Pfam
 13.1: 1313 multiple alignments and profile HMMs match the majority of proteins, Nucleic Acids Res., 27, 260-262.
- 14. Kikuno, R., Nagase, T., Suyama, M., Waki, M., Hirosawa, M., and Ohara, O. 2000, HUGE: a database for human large proteins identified in Kazusa cDNA sequencing project, Nucleic Acids Res., 28, 331-332.
- Nagase, T., Ishikawa, K.-I., Suyama, M. et al. 1998, Prediction of the coding sequences of unidentified human genes. XI. The complete sequences of 100 new cDNA clones from brain which code for large proteins in vitro, DNA Res., 5, 277-276.
- Qualmann, B., Roos, J., DiGregorio, P. J., and Kelly, R. B. 1999, Syndapin I, a synaptic dynamin-binding protein that associates with the neural Wiskott-Aldrich syndrome protein, Mol. Biol. Cell, 10, 501-513.
- Steplewski, A., Haas, S., Amini, S., and Khalili, K. 1995, Regulation of mouse myelin basic protein gene transcription by a sequence-specific single-stranded DNA-binding protein in vitro, Gene, 154, 215-218.
- Ichtchenko, K., Nguyen, T., and Sudhof, T. C. 1996, Structures, alternative splicing, and neurexin binding of multiple neuroligins, J. Biol. Chem., 271, 2676-2682.

